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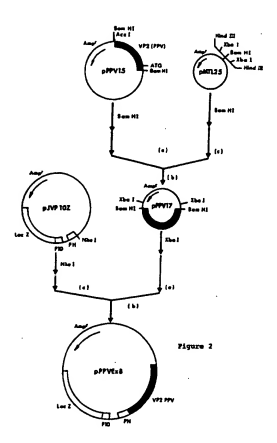
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METHOD FOR PRODUCING A SUBUNIT VACCINE AGAINST PORCINE PARVOVIRUS.

(57) Method for the production of a subunit vaccine against porcine parvovirus (PPV). The method is comprised of a first step wherein is obtained a recombinant protein VP2 of PPV by using the replication of a recombinant baculovirus wherein the gene corresponing to VP2 has been previously inserted in cells of a permissive host. The protein VP2 obtained in this invention has the capacity of forming mpty chimeric capsides with high immunogenicity and can be provided as vaccin formulations for protecting pigs against the infection caused by PPV. Additionally, they can be chemically or genetically engineered in order to incorporate epitopes corresponding to other viral proteins and they can be used in the formulation of polyvalent vaccines. The recombinant baculovirus AcMNPV.pPPVEx8 expresses the VP2 of PPV in conditions making possible the formation of pseudo-viral capsid s, and has been deposited at the ECACC. These vaccin s find application in the veterinary field.



FIELD OF THE INVENTION

The present inv ntion relat s in general to viral proteins and to assays and vaccines using the same and, in particular, to a protein related to the major antigen (VP2) of the Porcine Parvovirus (PPV) capsid. Such protein was produced in an expression vector of baculoviruses multiplied in a cell culture of a permissive host. The protein obtained with the instant invention is singularly characterized in forming empty chimeric capsids that are useful in vaccine formulation.

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BACKGROUND OF THE INVENTION

The Porcine Parvovirus (PPV) causes reproductive failure in swine, resulting in death and foetal mummification, still births and other reproductive failures in pregnant sows. (Joo & Johnson. 1976. Veterinary Bulletin 46, 653-660; Mengeling. 1978. J. Am. Vet. Med. Assoc. 172, 1291-1294). PPV is an autonomous parvovirus containing a single strand DNA molecule of approximately 5000 nucleotides (Mollitor et al. 1984, Virology 137, 241-254). The complete sequence of this genome has been recently described by our group (Ranz et al. 1989. J. Gen. Virol. 70, 2541-2553). Four virusspecific proteins have been described: three capsid proteins (VP1, VP2 and VP3 of Mr values 83000, 64000 and 60000 daltons, respectively) and one non structural protein NS1.

The PPV is related to the Kilham rat virus (KRV) group of autonomous parvoviruses formed by KRV, minute virus of mice (MVM), *LullI*, H-1, Feline Panleukopenia virus (FPV), canine parvovirus (CPV) and the mink enteritis virus (MEV). These viruses share several common features with other autonomous parvoviruses:

- 1. There are two large open reading frames (ORFs).
- 2. The mRNAs from both ORFs are polyadenylated and 3'-coterminal.
- The left ORF encodes non-capsid proteins which are necessary for viral DNA replication and the right ORF encodes the major capsid proteins as a nested set.

To date, there are several vaccines protecting from porcine parvovirus disease, which are based on conventional inactivation methods of the virus. However, every previous attempt of new vaccines production using recombinant proteins produced in procariotic microorganisms (v.g. *E.coli*) have failed. In this invention, a new process is described for obtaining a new kind of vaccines based on the immunogenic properties of the major protein VP2, expressed in a baculovirus system multiplied in a cell culture of a permissive host.

For the last years, our laboratory has been studying with great detail the molecular biology of PPV. The findings obtained thus are summarized in two pioneer publications:

- A. Ranz, J.J. Manclus, E. Díaz Aroca, J.I. Casal. 1989. Porcine Parvovirus: DNA sequence and genome organization. J. Gen. virol. 70, 2541-25463.
- J.I. Casal, E. Díaz, A. Ranz, J.J. Manclus. 1990. Construction of an infectious genomic clone of PPV. Effect of the 5' end on DNA replication. Virology 177, 764-767.

These publications are related with the knowledge of the viral DNA sequences encoding the proteins forming the viral capsid. These sequences allowed the identification of the gene that encodes the VP2 of PPV and its manipulation and insertion into the specific vectors to be expressed in the baculovirus system. This system allows a large-scale protein production based upon the replication of recombinant baculovirus derived from the Autographa Californica nuclear polyhedrosis virus (AcmnPV) in insect cell cultures. The state of the art for these vectors is summed up in two scientific papers as follows:

- Luckow, V.A. & Summers, M.D. 1988. Trends in the development of baculovirus expression vectors. Bio/Technology 6, 47-55.
- 2. Vialard et al. 1990. Synthesis of the membrane fusion and hemagglutinin proteins of Measles virus using a novel baculovirus vector containing the β -galactosidase gen. J. Virol. 64, 37-50.

The advantages of VP2 protein synthesis in a baculovirus vector are remarkable over the virus production in cell culture and subsequent purification, in the economic cost of the process and immunogenic antigen output. On the other hand, this invention avoids the sacrifice of animals to stablish primary cell cultures for virus replication, to keep viral reservoires and the usual hazard in virus handling, etc.

SUMMARY OF THE INVENTION

The present invencion puts forth a new process for producing a recombinant subunit vaccine to protect pigs from PPV. The new vaccine produced thus can contain:

- 1. The PPV VP2 protein produced in an expression vector of baculoviruses multiplied in a cell culture of a permissive host (this protein hereinafter optionally refered to as "VP2 hereof") or
- Empty chimeric capsids formed by assembly of the VP2 hereof.

Th VP2 protein hereof is singularly characterized in forming empty chimeric capsids, optionally

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incorporating other viral protein epitopes by genetic manipulation of the recombinant baculoviruses or chemical manipulation of the chimeric capsids.

The object of the invention is therefore a new process for obtaining a new improved subunit vaccine capable of protecting pigs from PPV infections. As aforesaid, the said vaccine can either contain the VP2 protein hereof or empty chimeric capsids formed by the said VP2 protein hereof, in as much as the said empty capsids have an enhanced hemagglutinant activity and are highly immunogenic, excelling other recombinant proteins of these viruses produced hereto in any other vector. The new vaccines the invention provides and being one of its objects can contain either the said empty capsids with an immunologically acceptable diluent, with or without an adjuvant, or the VP2 protein hereof together with a diluent and an adiuvant.

Since the said chimeric capsids can be chemically or genetically manipulated to introduce other unrelated viral protein or peptide epitopes therein, the use of the said capsids both for PPV vaccinal purposes and modified to incorporate other epitopes, thereby to provide a polyvalent vaccine, are further additional objects of this invention.

The VP2 protein obtained with the invention and the chimeric capsids it can form can be useful in diagnosis to detect the presence of PPV specific antibodies or to induce polyclonal or monoclonal antibodies capable of PPV detection. The use of the VP2 protein hereof and of the chimeric capsids it can form for the above purposes is a further object of the present invention.

An additional object of this invention is a recombinant baculovirus and the process for obtaining the same, capable of producing a PPV VP2 r combinant protein identical to the viral protein, as shown in antigenic reactivity assays and other biological functionality assays. The recombinant baculovirus was called AcMNPV.pPPVEx8 and filed on 2.3.91 with the European Collection of Animal Cell Cultures, (ECACC), at Porton Down, Salisbury, Wiltshire SP4 OJG (Great Britain), accession number V91030213.

A further object of the invention is the new baculovirus transfer vector (pPPVEx8) containing the nucleic acid sequence coding for the VP2 here-of. With a process known as homologous recombination with AcMNPV wild-type genome this new vector leads to the said AcMNPV.pPPVEx8 recombinant baculovirus.

This invention also provides the nucleic acid sequenc coding for the VP2 protein of the invention (Figure 1).

The empty PPV chimeric capsids formed by autoass mbly of the PPV recombinant VP2 proteins are yet another object of this invention.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1, shows the nucleotide s quenc coding for the VP2 hereof and the amino acid sequence thereof. The nucleotide sequence is shown in the direction 5'-3' from left to right. The amino acids have been designated using the generally accepted three-letter code.

Figure 2, shows the construction of the pPPVEx8 transfer vector, pointing out the appropriate manipulations for inserting the PPV VP2 gene in the pJVP10Z plasmid.

Figure 3, shows the presence of empty chimeric capsids formed by aggregation of the VP2 protein hereof, as observed under an electron microscope.

Figure 4, shows the antibodies titre average values of sera from pigs immunized two times with N3 μ g of empty PPV chimeric capsids adjuvanted with Alhydrogel + QuilA.

The antibody titre was measured by:

A) ELISA anti PPV virions



B) PPV hemaglutination inhibition assays (HI)



C) PPV Neutralization



Figure 5, shows the antibodies titre values against PPV obtained by ELISA in pregnant sows vaccinated with PPV chimeric capsids (▲,▼) and non-vaccinated (■), all of them were challenged with a PPV virulent strain.

45 DETAILED DESCRIPTION OF THE INVENTION

The invention provides a new process for obtaining a recombinant subunit vaccine serving to protect pigs against infections due to Porcine Parvovirus. The vaccine can contain the PPV VP2 recombinant protein produced in an expression vector of baculovirus multiplied in a cell culture of a lepidoptera or other permissive host, or chimeric capsids formed by aggregating the said r combinant VP2.

The invention also provides a recombinant baculovirus capabl of expr ssing the PPV VP2 when inoculated in a p missiv host, and th

process for obtaining the said recombinant baculovirus.

The obtention of the recombinant baculovirus basically comprises the following steps:

- a) Preparing the gene coding for the PPV VP2;
- b) inserting the VP2 gene in a baculovirus transfer vector:
- c) transfecting permissive host cells with the said baculovirus transfer vector holding the VP2 gene; and
- d) selecting the recombinant baculovirus expressing the PPV VP2 protein.

The recombinant baculovirus obtained thus and the proteins and capsids produced are additionally characterised. These steps will be described in detail hereinafter.

In a preferred embodiment the gene coding for the PPV VP2 protein is obtained from plasmid pPPV15, previously constructed in our laboratory, containing the sequences encoding for VP2, and inserted in the unique cloning site Nhel of the AcMNPV-derived pJVP10Z plasmid, thereby to obtain a baculovirus transfer vector. In our invention the pPPVEx8 vector proved to have the PPV DNA adequately oriented to be expressed by the AcMNPV virus polyhedrin promoter.

The pPPVEx8 vector was used to co-transfect permissive host cells, with the AcMNPV virus wild-type DNA. Reference could, inter alia, be made to cells of lepidoptera or their larvae. In a preferred embodiment of this invention *Spodoptera frugiperda* (S. frugiperda) cells, generally from the Sf9 strain, were transfected using pPPVEx8, though it can naturally be assumed that similar results would be achieved transfecting other permissive cells for recombinant baculovirus replication.

After transfection, the recombinant baculoviruses were selected after removing and titrating the supernatant produced in a confluent monolayer of *S.frugiperda* cells. The blue plates with no trace of the viral polyhedrin under a light microscope were collected and back-titrated on *S.frugiperda* cells to obtain the recombinant baculoviruses. The AcMNPV.pPPVEx8 recombinant baculovirus is capable of expressing the PPV VP2 recombinant protein (VP2 hereof) and was filed with the ECACC, accession number V91030213.

A Dot Blot assay was used to verify that the VP2 gene had been adequately integrated into the said recombinant baculovirus genome.

The proteins expressed by the *S.frugiperda* cells infected with the recombinant baculovirus were analysed by electrophoresis in 8% to 15% SDS-polyacrylamide gradient gels and were stained with Coomassie blue to observe the presnc of a protein with a virtual molecular weight of 64 KDa, equivalent to that of the viral VP2 in the

recombinant virus plate. Immunodetection assays showed that the anti-PPV polyclonal antis rar acted with the VP2 expressed by the r combinant baculovirus. It can in light of these results be said that the VP2 hereof expressed by the recombinant baculovirus in *S.frugiperda* cells is antigenically undistinguishable from the viral VP2.

The VP2 protein obtained with the above-described process can be used for diagnosis purposes to detect the presence of specific PPV antibodies or to induce polyclonal or monoclonal antibodies capable of PPV detection. They can further be used to immunise animals to PPV. ELISA assays have shown that immunised animal sera recognised the viral antigens while hemagglutination inhibition assays (HI) showed that sera from animals immunised with the purified VP2 protein obtained with this invention offered HI titres of 1/320 when 4HA units of purified PPV were used as antigen. Thus, it can be said that the animals immunized with the VP2 obtained with our process are highly protected.

Based on the results obtained the VP2 protein expressed by the recombinant baculovirus vector hereof can be used to be formulated in vaccines in order to protect pigs from infections caused by PPV. These vaccines can be both passive and active. A passive vaccine could be obtained immunising animals with the recombinant and purified VP2 hereof and then isolating polyclonal antibodies from the said VP2 which could when purified be used in therapeutic or prophylactic applications. An active vaccine can be prepared resuspending the VP2 hereof in an immunologically acceptable diluent with an adjuvant.

It was submitted above that the VP2 protein obtained with the process of this invention is peculiar in that it can be aggregated, working pursuant to our conditions, and form pseudo-viral empty chimeric capsids of regular and uniform structure and with a size of about 22 nm, as shown by electron microscopy. No-one had to date described the "in vitro" formation of pseudo-viral capsids in Porcine Parvovirus using only the VP2 protein thereof. This much allows the recombinant VP2 proteins obtained to be easily purified. Furthermore, the empty capsids formed by VP2 assembly have an enhanced hemagglutination activity and are highly immunogenic, more so than other PPV recombinant proteins produced heretofore in other vectors. The said capsids can hence be formulated to be used in vaccines capable of protecting animals from infections caused by PPV. Broadly speaking, an active vaccine can be prepared resuspending the said capsids in an immunologically acceptable diluent, with or without an adjuvant. An important feature of these empty chimeric capsids, that could be obvious to someone skilled in the art.

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is that they can be ch mically or genetically manipulated to introduce the protein epitopes of other viruses, infection of which is to b protected, thereby to work as a polyvalent vaccine.

Phosphate buffered saline (PBS) solutions or other saline-like solutions could be used as an immunologically acceptable diluent. The adjuvant used could be alumina gel suspensions or other adjuvants regularly used in formulating vaccines.

DETAILED DESCRIPTION OF A PREFERRED EMBODIMENT OF THE INVENTION (EXAMPLE)

1. OBTAINING RECOMBINANT BACULOVIRUSES EXPRESSING THE PPV VP2 GENE

1.1. Preparing the VP2 gene of PPV

The complete PPV genome was cloned in our laboratory for the first time in the bacterial plasmid pUC18, obtaining the genomic clone pPPV10. The construction of this genomic clone has been described in "Construction of an infectious genomic clone of Porcine Parvovirus: Effect of the 5'-end on DNA replication", Casal et al. 1990. Virology 177, 764-767. As starting material we employed as original virus the PPV strain NADL-2, available at the American Type Culture Collection, Rockville, Maryland (USA), accession number ATCC-VR742.

After manipulation, the pPPV10 genomic clone was used to obtain the pPPV15 plasmid (Figure 2), which contains all the coding sequences for PPV VP2 in a DNA fragment of approximately 1.9 Kbp. Th foreign DNA was inserted so as to allow its extraction in a single step, thanks to a BamHI restriction site. The inserted 1.9 Kbp DNA was isolated by electrophoresis in low melting agarose gels and inserted in a previously dephosphorilated and BamHI digested pMTL-25 vector.

The plasmid so obtained was called pPPV17. It had the VP2 coding sequences from the PPV virus flanked by two Xbal restriction sites.

1.2. VP2 gene insertion in a baculovirus transfer vector

The plasmidic vector with AcMNPV-derived Nhel site, (pJVP10Z plasmid), (Vialard, J. Et al. J. Virol. 64, 37-50, 1990), was a gift of Dr. Cris Richardson (NRC. Quebec. Canadá) and was used to clone the Xbal fragment obtained from the pPPV17 plasmid as described in Figure 2. As the said figure shows, the Xbal pPPV17 fragment containing the PPV VP2 coding sequence was inserted into the pJVP10Z Nhel site.

The plasmids thus obtained, containing the PPV VP2 gene, were purified according to the

alkaline lysis method (Birmboim & Doly. Nucl ic Acids R s. 7, 1513-1523. 1979) and charact riz d by r striction endonuclease mapping.

The pPPVEx8 recombinant plasmid proved to have the PPV VP2 gene placed in the correct orientation to allow its expression by the AcMNPV virus polyhedrin promoter.

1.3. Recombinant virus transfection and selection

S.frugiperda cells were transfected with infectious DNA mixtures of AcMNPV purified DNA and pPPVEx8 plasmidic DNA, according to the process described by Burand et al. Virology 101. 286-290. 1980. AcMNPV DNA (1 μg) purified according to Smith and Summers (Virology 123, 293-406. 1983), was mixed with two different quantities of plasmidic DNA (1 and 5 μg) and taken to 750 μl with a Hepes buffered saline solution (25 mM Hepes, pH 7.1, 140 mM NaCl and 125 mM CaCl₂). The DNA solution was inoculated on to monolayer cultures of 2x10⁶ S.frugiperda cells and incubated for 4h at room temperature. The supernatant was removed and 5 ml of medium containing 10% foetal calf serum were added.

After 4 days of incubation, supernatant was collected and plated on confluent monolayers of *S.frugiperda* cells. To improve the detection of recombinant plates, a X-gal blue indicator was added to the agarose. Blue plates showing no traces of occlusion bodies (viral polyhedrin) under a light microscope were collected and replated on *S.frugiperda* cells to obtain high-titre stocks of the recombinant viruses (10⁷⁻⁸ pfu/ml).

The recombinant baculovirus was called AcMNPV.pPPVEx8 and filed at the European Collection of Animal Cell Cultures (ECACC), accession number V91030213.

2. DOT BLOT ASSAY

A Dot Blot assay was made to determine whether the VP2 gene had been integrated in the recombinant baculovirus genome, as follows.

To obtain DNA from the recombinant baculovirus the *S.frugiperda* cells were infected with the said virus at a multiplicity of infection of 5 PFU/cell and incubated at 27 °C for 48 hours. The infected cells were collected, sonicated and centrifuged at 1000 rpm for 10 min to eliminate cell debris. The supernatant was used as starting material for the assays.

A volume of 100 μ l was denatured with 10 μ l of 1M NaOH, boiled for 5 min and placed immediately on ice. The mixture was neutralized with 10 μ l of 1M PO₄H₂Na. A 20xSSC solution was added immediately to obtain a final 6xSSC concentration

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(SSC, solution saline citrate).

The solution was transferred to a nitrocellulose filter that had previously been moistened with 6xSSC. It was washed with more 6xSSC and dried at 37 °C for 30 min. The DNA was fixed to the nitrocellulose filter under an u.v. light for 2-3 min. The membranes were then hybridized with a specific probe of the VP2 DNA region labeled with Phosphorous-32 at 37 °C overnight. It was afterwards washed with decreasing SSC solutions and autoradiographed.

A strong sign of hybridization was observed only in those wells containing supernatants from the cultures infected with recombinant viruses, which indicated that the VP2 gene had been integrated in the viral genome.

3. PROTEIN AND IMMUNODETECTION ANALYSIS

S.frugiperda cells were infected with the recombinant baculovirus at a multiplicity of 5 PFU/cell and incubated at 27 °C for 48 h. The cells were collected by centrifugation at 1000 rpm for 10 min, washed twice with phosphate buffered saline (PBS) solution pH 7.4 and resuspended at 1x10⁶ cells/ml in lysis buffer (5% sodium dodecyl sulphate (SDS), 1% β -mercaptoethanol and 17.4% glycerol). The samples were charged in 8 to 15% SDS-polyacrylamide gradient gels for electrophoresis and stained with Coomassie blue or transferred to nitrocellulose membranes for immunodetection. The Coomassie blue showed the major presence of a protein with a molecular weight of 67 KDa, equivalent to that of the viral VP2, in the recombinant virus lane.

For immunodetection the proteins were transferred to nitrocellulose membranes in accordance with previously described methods (Burnette, Anal. Biochem. 112. 195-203, 1981. Towbin eta al., Proc. Natl. Acad. Sci. USA 76. 4350-4354. 1979). Protein transfer was made in a PhastSystem (Pharmacy) apparatus. In general 25 mA/gel were used for 10-15 min. The nitrocellulose strips were blocked with 3% powdered skim milk in 20 mM Tris HCl pH 7.5. 500 mM NaCl (TBS) for 30 min at room temperature. The strips were then incubated for an hour at room temperature with the first anti-PPV rabbit aniserum, washed with TBS-0.05% Tween-20 for 30 min at room temperature and incubated for 1 h at room temperature with goat anti-rabbit serum labeled with biotin (1:500). The strips were washed again and allowed to react with streptavidin labeled with peroxidase (1:2000) for 30 min at room temperature. After a thorough wash, the filters were developed with a TBS solution containing 0.5 mg/ml of 4-chloro-1-naphthol (Sigma), 17% (v/v) of methanol and 0.015% of hydrogen peroxide in TBS

until visible bands appeared. The reaction was stopped rinsing the strips with distilled water.

All the rabbit policional antisera that had b en prepared against full PPV viral particles, reacted to the VP2 protein expressed by the recombinant baculovirus.

3.1. Recombinant protein and capsid purification

S.frugiperda cells were infected AcMNPV.pPPVEx8 recombinant virus at a multiplicity of infection of 5-10 PFU/cell and incubated at 27°C for 48-72 h. The cells were collected by centrifugation at 1000 rpm for 10 min, washed twice in a phosphate buffered saline solution pH 7.4 and resuspended at 2×10^7 cells/ml in a 25 mM bicarbonate buffer pH 9.5. The resuspended cells were broken down by sonication and centrifuged at 10.000 rpm for 10 min to eliminate cell debris. Supernatant containing the recombinant VP2 protein can be purified using its autoaggregating capacity to form empty capsids. They are to this end either purified by 20% ammonium sulphate precipitation or the empty capsids are centrifuged on CsCl gradients at 45,000 rpm for 14 h. Capsids offer a buoyant density (p) of 1.30 g/cm3 when banded in CsCl gradient. Preparation purity was determined by electrophoresis in SDSpolyacrylamide gels as the method described above and VP2 protein purity turned out to be in excess of 99%.

4. HEMAGGLUTINATION ACTIVITY

Hemagglutination activity of the VP2 protein was assayed in accordance with an already known method (Joo, H.S. et al. Aust. Vet. J. 52: 422-424. 1976). This functional activity is exclusively associated to the particulate character of the product, thereby clearly distinguishing the same from previous ones.

The results show that the preparation of VP2 capsids has an hemagglutination titre of 5.10⁵ units/ml.

5. CONFIRMING THE PRESENCE OF CAPSIDS BY ELECTRON MICROSCOPY

A purified VP2 preparation was stained by negative contrast with 2% uranyl acetate and observed under an electron microscope at a magnifying powr of 40.000 x 2.5, the pr sence of a large number of pseudo-viral chim ric particles of regular and uniform structure and with a size of roughly 22 nm was observed. (Figure 3).

6. IMMUNIZING RABBITS

Two New Zealand rabbits weighing 2 Kg wer immunized three times intramuscularly with 100 µg of a purified VP2 preparation. The first time in a complete Freund adjuvant, the second and third times with an incomplete adjuvant. A week after the last immunization, the rabbits were bled and the sera obtained tested with an ELISA assay and an hemagglutination inhibition assay, as described hereinbelow.

6.1. Quantification of the anti PPV antibodies by ELISA and HI

The presence of PPV specific antibodies in the serum of immunized animals was determined by means of an indirect ELISA assay. The antigen used was both purified PPV virus and purified VP2 protein. Briefly, polystyrene plates were coated with 0.5 µg of virus or 0.25 µg/well of VP2 in 100 μl of carbonate buffer (0.05 M, pH 9.6) at 4°C overnight. The plates were washed with PBS (0.15 M NaCl in 0.1 M sodium phosphate pH 7.4) containing 0.05% Tween-20 and incubated with the anti-PPV rabbit antiserum for 2 h at 37 °C, washed again and incubated with goat anti-rabbit IgG biotin-labeled. Then, streptavidin labeled with peroxidase was added to the antibody marked with biotin for 30 min at room temperature. The plates were washed again and the reaction developed with o-phenylenediamine as a substrate for the peroxidase, for 10 min in darkness, and read at 450 nm in a multichannel spectrophotometer.

For the rabbit sera, the ELISA titre achieved was 1/1600 against the VP2 protein and 1/3200 against the PPV complete virus.

The HI assay was made according to the standard methods previously described (Joo, H.S. et al Aust. Vet. J. 52: 422-424. 1976). The titre obtained was 1/320 when 4 HA units of purified PPV were us d as antigen. Based on the good correlation observed between the HI and the protection titres of the animals against virus infection, we assume that the animals able to develop this titre have a high protection level.

7. PIG IMMUNIZATION

Two pigs were immunized with purified preparations of PPV VP2 chimeric capsids made of recombinant autoassembled VP2 proteins. One of the pigs was completely seronegativ while the other one had a low level of antibodies, due to the fact that his mother had been previously vaccinated. Both pigs were kept in an isolation unit during the assay.

The vaccine was formulated by mixing the capsids (N3 µg) with a standard adjuvant syst m: 50% Alhydrogel (Superfos. Denmark) + 500 μg QuilA (Superfos). All components were kept in standard concentrations except for the inactivated virus that was changed by PPV capsids. The pigs received two subcutaneous vaccinations: the first time the dose was 2 ml, and three weeks afterwards they were injected with a second dose of 1 ml. The serum was tested weekly, before vaccination and up to ten days after the last dose. The presence of anti-PPV antibodies in the pig serum was determined according to three different methods: (1) ELISA anti-PPV virion assay (2) Haemagglutination inhibition assay (Joo et al. 1976. Aust. Vet. J. 52: 422-424) and (3) PPV neutralization assay (Holm-Jensen M. 1991. Acta Vet. Scand. 22: 85-98).

The titres obtained with the capsids recombinant vaccine were similar (Figure 4) to those normally obtained with the inactivated commercial vaccine. These results show that the PPV chimeric capsids are highly immunogenic and able to replace the inactivated virions. Even although one of the pigs kept residual maternal antibodies, the response to the vaccine was not inhibited, reaching levels similar to those obtained in the seronegative pig.

The response was achieved with a low dose of the recombinant capsids ($N3~\mu g$), which shows that the product shows high prospects of commercial application as vaccine.

8. "In vivo" PROTECTION AGAINST A VIRU-LENT VIRUS

A challenge experiment was carried out to study the suitability of the recombinant VP2 capsids in the induction of protective immunity against PPV in pregnant sows.

Before artificial insemination, 2 seronegative sows were vaccinated with the same vaccine preparation described in the previous example. The antigen content, adyuvants and formulation of the vaccine were the same. Both sows were vaccinated twice with a 3 week interval. A seronegative pregnant sow was used as non-vaccinated control. At about 40 days of gestation the 3 pregnant sows were inoculated by intravenously route with 107 TCDIso of a virulent PPV strain "839" (Sørensen and Askaa. 1981. Acta Vet. Scand. 22, 171-179). The three sows were sacrificed at 66 days of gestation.

Crown to rump length and gross pathological lesions (GPL) were recorded for each foetus. Blood samples of umbilical cord and sera from the foetuses wer collected and checked for th presence of anti-PPV specific antibodies by an indirect

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immunofluorescence test (IFAT) (Sørensen et al. 1980, Acta Vet. Scand. 21, 312-317). Anti-PPV anti-bodies wer also checked by a t st of countercurrent immunoelectrophoresis. Beside this, these samples were checked also for IgM and IgG content using rocket electrophoresis (Dalsgaard et al. 1979. Acta Vet. Scand. 20, 312-320). In those cases where no blood from umbilical cord could be obtained, abdominal fluids or brain tissue extracts were used.

Foetal kidney, liver and lung tissues were collected and analyzed for the presence of PPV antigen by the ELISA test, routinely used in the State Veterinary Institute for Virus Research, Lindholm (SVIV) for PPV diagnosis. Also, blood samples of the sows were taken before vaccination, in the revaccination, 10 days after, in the time of the viral inoculation and in the sacrifice and checked for the presence of anti-PPV antibodies by the ELISA test previously mentioned.

The three sows remained healthy throughout the experiment. The antibody titers obtained during the experiment are shown in Figure 5. The non-vaccinated sow remained seronegative until the viral challenge. After infection a dramatic increase in the anti-PPV antibody titers was observed in the necropsy. The two vaccinated sows show antibody titers that increase after vaccination and revaccination. These titers suffer a posterior slight increase due to the administration of virulent virus.

a) Foetuses of the non-vaccinated control

At necropsy, foetuses from the sow #1451 (non-vaccinated control) displayed typical lesions of intrauterine PPV infection (Bachmann et al. 1975, Infect Immunity, 12, 455-460; Joo et al. 1976; Arch. Virol. 51, 123-391; Nielsen et al. 1991. Vet. Microbiol. 28, 1-11). Four foetuses were alive. One of them did not show GPL, however, the other three displayed GPL of varying severity, typically discoloration, morbidity, with large volumes of ascitic fluids, edema, pulmonary stasis and erythema, thymic atrophy and enlargement of the liver. Other five foetuses were dead and had severe GPL including growth retardation. Universal extreme edema, hyperemia and pronounced tissue destruction. Three mummified foetuses had CR lengths of 11.5 to 12.5 cm indicating growth arrest at 57 days of gestation.

PPV antigen was detected in all the foetuses of the non-vaccinated sow, using the ELISA test previously described. In pleural fluids of three foetuses was detected antibody r sponse (foetal) anti-PPV to the virus used for the challenge, as measured by IFAT and counter-current immunoelectrofor sis. The presence in these samples of IgG and IgM was confirmed by "rocket" immunoelectrophoresis.

The pig foetuses are able to induc an anti-PPV antibodies at 60 days of gestation (J. Niels n et al. 1991, Vet. Microbiol. 28, 1-11).

b) Foetuses of the vaccinated sows

In the vaccinated sows, one of them had 10 foetuses and the other 8 foetuses. All of them were alive and normal in the necrospsy. All of them appeared healthy. No PPV antigen was detected in any of the foetuses. Neither anti-PPV antibodies were detected in blood or pleural fluid by any of the used techniques. The absence of IgG or IgM immunoglobulines was confirmed by rocket immunoelectrophoresis.

On the basis of these results above described it is possible to conclude that recombinant VP2 capsids of PPV expressed in the baculovirus/insect cells system are able to induce a protective immunity against an intravenous innoculation with virulent PPV virus in pregnant sows.

Based on these results, it is demonstrated that recombinant VP2 capsids may constitute the base for a new range of commercial vaccines useful on the control of PPV infection in pigs. In the same way, since the essential immunodominant epitopes of PPV are expressed on the VP2 capsids, these capsids can be useful as a reagent in the diagnosis of PPV infection in pigs, for instance, in kits for antibodies detection.

9. VACCINE FORMULATION AGAINST THE IN-FECTION CAUSED BY PPV

It is possible to obtain a pasive vaccine immunizing animals with the recombinant VP2 capsids, purified as described in the present invention. Polyclonal antibodies, directed against this VP2, can be isolated from serum, milk or other animal bodily fluids. These antibodies can then be purified and used for therapeutics or prophylactic applications.

An active vaccine can be prepared resuspending the recombinant VP2 capsids described herein in an immunologically acceptable diluent such as PBS and an adjuvant such as Alhydrogel or QuilA. Initial and record injections or oral administration of the vaccinal solution can be used to confer immunity.

An active vaccine can also be prepared resuspending the empty capsids in an immunologically acceptable diluent with or without an adjuvant. Anyone skilled in the art will clearly see that these chimeric capsids formed only by VP2 can be chemically or genetically manipulated to introduce other viral protein epitopes and hence function as a polyvalent vaccin.

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10. CONCLUSIONS

The AcMNPV.pPPVEx8 baculovirus is capable of producing a recombinant VP2 absolutely identical to the viral VP2 protein, as shown with the DNA sequence, molecular weight estimate and antigenic characterization. The VP2 obtained herein with our process is also remarkably capable of forming empty capsids, thereby providing the same with hemagglutination and immunogenic activity that are clearly greater than in other previously described recombinant proteins, as shown with the animal immunization tests herein described.

This enhanced immunogenic capacity can be used by those skilled in the art to present other viral protein epitopes, that can be introduced therein by either chemical or genetic manipulation of the recombinant baculoviruses.

Translation of the legends to the figures

Figure 2

- (a) Isolate agarose gel fragments.
- (b) Ligate.
- (c) Phosphatase

Figure 4

- (a) Logarithm of the titer.
- (b) Days post-immunization.
- (c) Second immunization.

Figure 5

- (a) Vaccination of pregnant sows with PPV chimeric capsids formed by recombinant VP2 autoassembly.
- (b) ELISA, antibody titres against PPV.
- (c) Days post-vaccination.
- (d) First vaccine.
- (e) Second vaccine.
- (f) Challenge.
- (g) Artificial insemination.
- (h) Necropsy.

The recombinant baculovirus has been filed with the European Collection of Animal Cell Cultures (ECACC) on March 4th, 1991.

Having described the object of the present invention what is deemed as the essence thereof is now contained in the following

Claims

- 1. A recombinant subunit vaccine to protect pigs against PPV infection, comprising:
 - a) an immunising quantity of:

- i) recombinant VP2 protein of PPV, or alternatively of
- ii) empty chimeric capsids form d by autoassembly of PPV recombinant VP2 protein; and
- b) a diluent and, optionally, an adjuvant, being immunologically acceptable.
- 2. A vaccine as in claim 1, characterized in that the said recombinant VP2 protein has been obtained by culture of permissive insect cells infected with a recombinant baculovirus with the gene coding for the PPV recombinant VP2 protein integrated in its genome.
 - 3. A vaccine as in claim 1, characterised in that the said empty chimeric capsids have been obtained by the autoassembly of the PPV recombinant VP2 proteins produced during replication in permissive insect cells of a recombinant baculovirus with the gene coding for the PPV recombinant VP2 protein integrated in its genome.
- 4. A vaccine as in claims 2 and 3, characterised in that the said recombinant baculovirus has been called AcMNPV.pPPVEx8 and filed with ECACC, accession number V91030213.
- 30 5. A polyvalent vaccine to protect animals from PPV, comprising:
 - a) an immunizing quantity of empty chimeric capsids formed by autoassembly of CPV recombinant VP2 proteins, chemically manipulated to introduce therein peptide or protein epitopes of viruses in respect of which infection is to be protected; and
 - b) a diluent and, optionally, an adjuvant, being immunologically acceptable.
 - 6. A vaccine as in claim 5, characterized in that the said empty chimeric capsids have been obtained by the autoassembly of the PPV recombinant VP2 proteins produced during replication in permissive insect cells of a recombinant baculovirus with the gene coding for the PPV recombinant VP2 protein integrated in its genome.
 - A vaccine as in claim 6, characterized in that the said recombinant baculovirus has been called AcMNPV.pPPVEx8 and filed with EC-ACC, accession number V91030213.
- 55 8. A polyvalent vaccine to protect animals from PPV and other viruses, comprising:
 - a) an immunizing quantity of mpty chimeric capsids formed by autoassembly

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of PPV recombinant VP2 proteins moreover containing the peptide or protein epitopes of viruses in respect of which infection is to be protected; and

- b) a diluent and, optionally, an adjuvant, being immunologically acceptable.
- 9. A vaccine as in claim 8, characterized in that the said empty chimeric capsids containing viral protein or peptide epitopes have been obtained by:
 - a) genetic manipulation of a recombinant baculovirus with the gene coding for the PPV VP2 in which the said epitopes have been introduced and integrated in its genome;
 - b) infection of permissive insect cells with the said genetically manipulated recombinant baculovirus; and
 - c) culture of the infected cells in conditions allowing the genetically manipulated VP2 chimeric capsids incorporating the relevant viral epitopes to be produced.
- 10. A vaccine as in claim 9, characterized in that the said recombinant baculovirus susceptible of genetic manipulation to introduce viral epitopes has been called AcMNPV.pPVEx8 and filed with ECACC, accession number V91030213.
- 11. A PPV recombinant VP2 protein comprising an amino acid sequence substantially identical to that shown in Figure 1 and that is antigenically identical to the PPV natural VP2 protein.
- 12. A protein as in claim 11 characterized in that has been obtained by culture of permissive insect cells infected with a recombinant baculovirus with the gene coding for the PPV recombinant VP2 protein integrated in its genome.
- 13. A protein as in claim 11, characterized in that, in addition, shows the capability to form empty chimeric capsids by autoassembly.
- 14. A protein as in claim 12 characterized in that the said recombinant baculovirus has been called AcMNPV.pPPVEx8 and filed with EC-ACC, accession number V91030213.
- 15. Empty PPV chimeric capsids characterized by having been obtained by autoassembly of PPV recombinant VP2 proteins expr ssed in permissiv insect cells inf cted with a recombinant baculovirus having the gene coding for the PPV VP2 protein.

- 16. Capsids as in claim 15, charact rized by having an enhanced hemagglutination capacity and being highly immunogenic and thus b ing appropriate to be used in formulating subunit and polyvalent vaccines.
- 17. Capsids as in claim 15, characterized in that the said recombinant baculovirus capable of expressing PPV recombinant VP2 proteins has been called AcMNPV.pPPVEx8 and filed with ECACC, accession number V91030213.
- 18. Capsids as in claim 15, characterized in that contain all the PPV essential immunodominant epitopes being, therefore, adecuated for its use in the preparation of kits for anti-PPV antibodies diagnosis.
- 19. A recombinant baculovirus able to express PPV recombinant VP2 proteins in permissive insect cells, that are able to form empty chimeric recombinant capsids by autoassembly.
- 20. A recombinant DNA sequence which codes for the PPV recombinant VP2 protein and contains a nucleotide sequence substantially identical to that shown in Figure 1.

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Pigure l

60 ACA Thr	120 GGT G1y	180 GTT Val	240 TAC	300 GAT ABP	360 TGG Trp	420 AGT Ser	480 CCA Pro	540 AAT Abn
GCA	GTT Val	TTG	ACA Thr	GAC	GTG Val	GTT	TCA	ACC
TCT Ser	GGG Gly	66c 61y	GAA Glu	GAA Gln	GGA Gly	15 T	ACC	GAC Asp
TTG Leu	666 61y	GA3 Glu	CAC H18	GTA	TGG Trp	AAC Asn	GCA	CTA Leu
GAA Glu	GCT	666 61y	GAA Glu	ATG Met	GCA	ATA Ile	TCA	GCA
ACT Thr	GGT Gly	TTG	CC.	CAA Gln	AAC ABn	GAA Glu	GAA Glu	GTC Val
GCG Ala	AGG Arg	TAC	ATG	GGA Gly	GCT	ACA	ACA	ATG Met
GCA	GGT Gly	CAA Gln	AAT Aen	GCG	gat Abd	ATG	ATT Ile	TTA Leu
AAT Asn	666 61y	TTT Phe	CTA	TCG	ATA Ile	AAC Asn	ACA Thr	AGC
ATT Ile	660 61y	GAA Glu	CAT Hib	666 61y	CTA	AAC Aen	AAA Lys	GCA
30 CCT Pro	90 GGT Gly	150 ACA Thr	210 ATA Ile	270 TCA Ser	330 TCA Ser	390 TCC Ser	450 CTT Leu	S10 ACT Thr
AAC	GGC Gly	CAA Gln	CTC	GAA Glu	TGG Trp	ATA Ile	GTA	CTA Leu
CAC His	GGC Gly	AAT Abn	AGA Arg	TCA	CCT	TTA	GTA	gat Abp
CAA	666 61y	AAT Abn	TCA	AAT Asn	ACA	Glag	AAT Asn	AAT Abn
GAA Glu	GGT Gly	TTC	GCA	CTA	GTA	166 Trp	TTC	AAT Abn
GTG Val	666 61y	AGT Ser	CAC	GTA Val	ATG	GAC	ATA Ile	tat Tye
AAT	TCT Ser	GGT Gly	GCA	CAT	CAA Gln	GCG	GCA	ATA Ile
GAA Glu	GAA	ACA	ACT	ATA Ile	ACA	S C S	CAA Gln	AAA Lys
AGT Ser	AAT	TCT Ser	ATC Ile	AGA	CAC	AAT Asn	GAA Glu	Acc
ATG Met	GGA	GTG Val	AGA Arg	AAA Lys	GCA	TTC	rrr Phe	CC. Pro

PPVVP2 5'---3'

Figure 1 (cont'd)

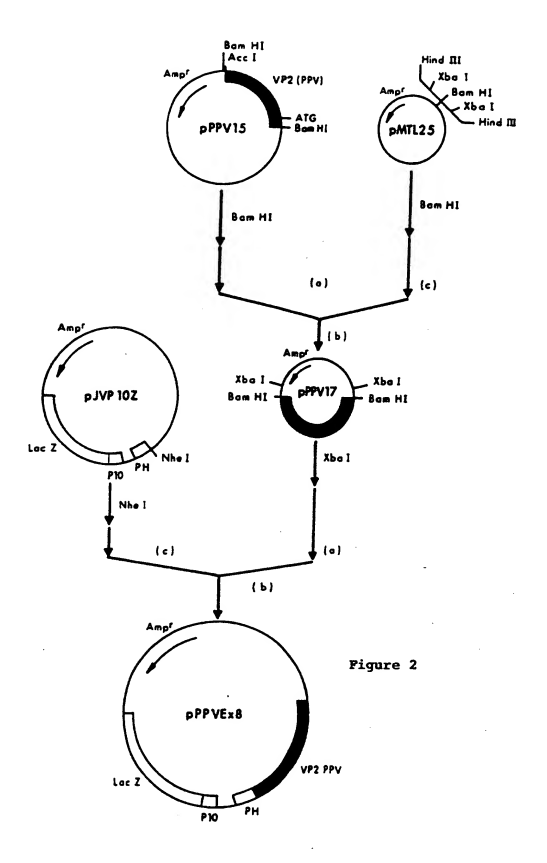
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CCA	AAT Asn	CAC	GGA Gly	TCA	GAA Glv	ACA	ACA Thr
TAT Tyr	CTA	CTA Leu	ACA	CAC H18	ACA Thr	S Lu	AAT Aen
TTT Phe	AAC Abn	AGG Arg	AGA Arg	ACT	ACC	CAC H18	TAT
GGT	AGA	AAC Asn	CTA Leu	TTA	CCT Pro	TAT	GGA Gly
CTT	ATC Ile	ACA	CTT Leu	AAA Lys	GAA Glu	GGT Gly	
ACA	TGC Cys	AAT Aen	CAT His	CTA	ACT Thr	AAA Lyb	CAG Gln
GAA Glu	TCA	CTC	ATT Ile	CCA	CTA Leu	AGA Arg	GCT
AGT	CTA	AGA Arg	CCA Pro	AAA Lys	CTA Leu	ACA Thr	CCA Pro
AGA	TAC	AAC	GTA Val	ACA	AAA Lys	AAC Asn	AGG
570 CCT Pro	630 TAT Tyr	690 AAT ABN	750 GCA Ala	810 GAC ABP	870 CCA Pro	930 GCT Ala	990 ATT Ile
GCA Ala	AGA Arg	CCA	AAT Asn	TTT Phe	CCT	GCA	GCA
GCA Ala	TAC	CAA Gln	GAA Glu	CAC	CTG	CCA	ACA
CCA	CAA Gln	TCA	ATA Ile	TAT Tyf	GGA Gly	CTA	GCA
ACA Thr	ACT	CAA Gln	ACA	ATA Ile	CTA	ACA	GAA Glu
TAC Tyr	CCA Pro	GGA Gly	TAC	gga gly	TCT Ser	GGA Gly	ACA Thr
CCA	aaa Lys	ACT Thr	TTC	ACA	aga	CCA	TAC
CTT	ACA	TAC Tyr	ATG	TCC Ser	AAC Asn	CAC	AGC
ACA	CCT Pro	ACA	ATT Ile	TTC	ACA	GAA Gla	AAT Aen
AAC	TTA	CCA	GAC	GAA Glu	Gla Gla	GAC	AAT

Pigure 1 (cont'd)

				-				
1080 GCA Ala	1140 CAA	AGT Ser	1260 AAT Asn	1320 TTC Phe	1380 TTT Phe	1440 GTT Val	1500 CCA Pro	1560 TCA Ser
ACA Thr	TAC Tyr	CAA Gln	GAA Glu	CAT H18	GTA Val	CAT CAT His	1500 GCA CCA Ala Pro	1560 TAT TCA Tyr Ser
CCA Pro	GAT Asp	CCA	CTA	ATG Met	CCT	CTA	ATA Ile	ACT
GTA Val	ATG Met	AAT Aen	AAC Asn	AAC Asn	GCA	AGA	AAA Lys	ATA Ile
ATA Ile	ACA	TTC Phe	CTA	TCT	ACT Thr	CCT	GTA Val	ATA Ile
CCT	TTT Phe	ACA	CCA	AAA Lyb	AAT Abn	AAA Lys	TTT Phe	AGA Arg
ACT	AGA TTT Arg Phe	TAC	GCA	666 61y	CTA AAC AAT Leu Asn Asn	CTA	CTA Leu	CCT AGA ATA ATA Pro Arg Ile Ile
CTA Leu	ATA Ile	AGA	CAG Gln	GGA Gly	CTA Leu	gat Asp	CAA Gln	CAA Gln
TTT Phe	GCT	GAA Glu	CAA Gln	ATA Ile	GCA	ACA	GGA G1y	CAA Gln
CCA	GGT Gly	CTA	AAT Aen	CCA Pro	ACA	gat Asp	CCA	CCT Pro
1050 GGT GGA Gly Gly	1110 CCA AAT Pro Asn	1170 CAA GAG Gln Glu	1230 CAA TTT Gln Phe	1290 TCA GAT Ser Asp	1350 CCA TTA Pro Leu	1410 GAA CTT Glu Leu	1470 AAT CCA ABn Pro	1530 GAC TCT ASP Ser
GGT Gly	CCA Pro	GIN GIN	CAA Gln	TCA Ser	CCA Pro	GAA Glu	AAT Aen	GAC Asp
AAT Asn	GAA Glu	TCA	CAA Gln	CCT	GGA Gly	AAA Lys	AAC	GCT
TCC	gat Asp	TCT Ser	AAG Lys	TTA	TAT Tyr	GAT	AAA Lys	AAT Asn
TAC	gat Asp	ACA Thr	CCA Pro	CIT	ACA	TGG	TGT Cys	TTC
GAA Glu	AAT Asn	ACC	GCT	ACA	AAT	ATA Ile	GTT Val	gat Abd
TTT Phe	TAT Tyr	TTA	AGA	GGA Gly	CTC	Gla	TTT Phe	gat Asp
AAT Asn	GAA Gln	CAC His	GGA	AAT	ACA	GGT Gly	CCA	ACA
ATG Met	ACA	GGA G	TGT Cy 8	AAT Aen	AAT Asn	AAT Asn	GCT	CTA
TAC	GAC	CAT	AAA Lys	ACA	ATG Met	CCA	ACA	AAC Asn

Pigure 1 (cont'd)

1620 TGG Trp	1680 AAT Asn	1740 TAG End
ATG Met	ACA Thr	TAC TYT
AAT Asn	CCT	TTA
AGT Ser	ATT Ile	AAA Lys
TCC Ser	TAT Tyr	AGA Arg
AGA Arg	AAA Lys	CCA
ATG Met	GGT Gly	ATA Ile
AAA Lys	ATT	CTT
GCA Ala	AAC	Gln
ACA	GAA Glu	TCA
1590 TTC Phe	1650 GCA Ala	1710 1 TAT 1 TYF
ACA Thr	ACA Thr	GAA Glu
CTA	ACA Thr	CCA Pro
ACA	ACA Thr	TTT Phe
GGA Gly	CAC	ATG Met
AAA Lys	CAA Gln	AAA
TGG Trp	CAA Gln	ATA Ile
TGG Trp	ATT Ile	66c 61y
TTT Phe	CCT	GGT Gly
AAC Asn	AAC Asn	ATT 11



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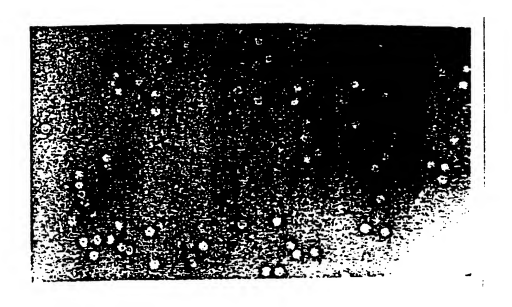


Figure 3

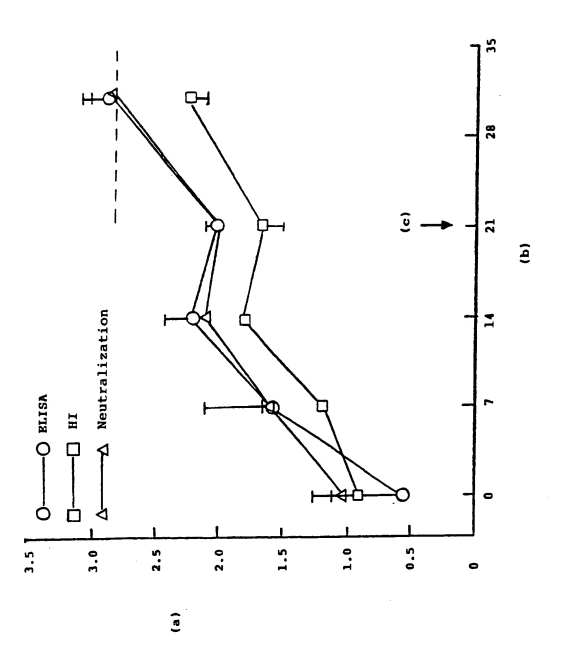
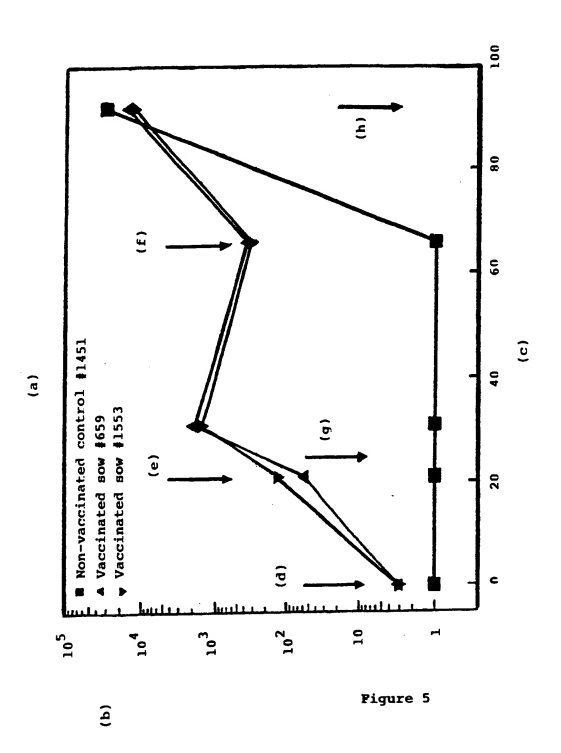


Figure 4



INTERNATIONAL SEARCH REPORT

International application No.
PCT/ES 92/00032

A. CLASSIFICATION OF SUBJECT MATTER Int.CI. 5 C 12 N 15/35 C 07 K 13/00 A 61 K 39/23 C 12 N 15/86						
According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED						
	ocumentation searched (classification system followed b	y classification symbols)				
In	t.Cl. ⁵ A 61 K C 07 K C 12 M	.				
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)						
C. DOCU	MENTS CONSIDERED TO BE RELEVANT					
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.			
X	Journal of General Virology, SGM (GB) A.I. Ranz et al.: " DNA sequence and genome orgar 2541-2553, see pages 2546-254 (cited in the application)	'Porcine parvovirus: nization", pages	11-14, 20			
Y	<u></u>		19			
γ .	EP, A, 0341611 (BOYCE THOMPSO PLANT RESEARCH, INC:) 15 Nove see the whole document	ON INSTITUTE FOR ember 1989,	19			
x	 WO, A, 8802026 (APPLIED BIOTE	ECHNOLOGY)	1-4,15-			
	24 March 1988, see the whole in particular pages 67-74; of	document	18			
Y			5–10			
						
		-/-				
Furthe	r documents are listed in the continuation of Box C.	See patent family amex.				
"A" docume:	categories of cited documents: nt defining the general state of the art which is not considered particular relevance	"T" later document published offer the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention				
"E" earlier d "L" document cited to	claimed invention cannot be cred to involve an inventive c					
"O" documer means						
"P" document the prior	P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family					
	Date of the actual completion of the international search 18 June 1992 (18.06.92) Date of mailing of the international search 26 August 1992 (26.08.92)					
Name and m	ailing address of the ISA/	Authorized officer				
EUROP	EUROPEAN PATENT OFFICE					
Facsimile No	3.	Telephone No.				

INTERNATIONAL SEARCH REPORT

International application No.
PCT/ES 92/00032

C (Continuat	ion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant p	assa ges	Relevant to claim No.
Y	DD, A, 286820 (HUMBOLDT-UNIVERSITAT BERLIN) 7 February 1991, see the whole document		5-10
x	EP, A, 0117767 (MOLECULAR GENETICS, INC.) 5 September 1984, see pages 90-110; claims 30 y 99-101		1-4
A	Journal of Virology, Vol. 64, No. 1, January American Soc. for Microbiology (US), J. Vialard et al.: "Synthesis of the membran fusion and hemagglutinin proteins of measles using a novel baculovirus vector containing t beta-galactosidase gene", pages 37-50; see the whole document	e virus,	1–20
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